

# **Responders and non-responders to influenza vaccination: a DNA methylation approach on blood cells**

## **Authors:**

Noémie Gensous <sup>a</sup>

Claudio Franceschi <sup>a,b,c</sup>

Bonnie B. Blomberg <sup>g</sup>

Chiara Pirazzini <sup>c</sup>

Francesco Ravaoli <sup>a</sup>

Davide Gentilini <sup>d</sup>

Anna Maria Di Blasio <sup>d</sup>

Paolo Garagnani <sup>a,b,c,f,g,h</sup>

Daniela Frasca <sup>i,\*</sup>

Maria Giulia Bacalini <sup>c,\*</sup>

\* Co-senior authorship

a: Department of Experimental, Diagnostic and Specialty Medicine, University of Bologna, Bologna, Italy.

b: Interdepartmental Center “L. Galvani,” University of Bologna, Bologna, Italy.

c: IRCCS Institute of Neurological Sciences, Bologna, Italy.

d: Istituto Auxologico Italiano IRCCS, Cusano Milanino, Milan, Italy.

e: Center for Applied Biomedical Research (CRBA), St. Orsola-Malpighi University Hospital, Bologna, Italy.

f: Clinical Chemistry, Department of Laboratory Medicine, Karolinska Institutet at Huddinge University Hospital, S-141 86 Stockholm, Sweden.

g: Institute of Molecular Genetics (IGM)-CNR, Unit of Bologna, Bologna, Italy

h: Laboratory of Musculoskeletal Cell Biology, Rizzoli Orthopaedic Institute, Bologna, Italy.

i: Department of Microbiology and Immunology, University of Miami Miller School of Medicine, Miami FL, USA.

## **E-mail addresses:**

Noémie Gensous: noemie\_gensous@hotmail.com

Claudio Franceschi: claudio.franceschi@unibo.it

Bonnie B. Blomberg: bblomber@med.miami.edu

Chiara Pirazzini: chiara.pirazzini5@unibo.it

Francesco Ravaoli: francesco.ravaoli2@unibo.it

Davide Gentilini: gentilini.davide@gmail.com

Anna Maria Di Blasio: a.diblasio@auxologico.it

Paolo Garagnani: paolo.garagnani2@unibo.it

Daniela Frasca: dfrasca@med.miami.edu

Maria Giulia Bacalini: mariagiuli.bacalini2@unibo.it

**Corresponding author:**

Paolo Garagnani

Alma Mater Studiorum – University of Bologna

Department of Experimental, Diagnostic and Specialty Medicine

Via San Giacomo 12

40126 Bologna (BO)

Tel: +39 0512094776

Mail: [paolo.garagnani2@unibo.it](mailto:paolo.garagnani2@unibo.it)

## Abstract

Several evidences indicate that aging negatively affects the effectiveness of influenza vaccination. Although it is well established that immunosenescence has an important role in vaccination response, the molecular pathways underlying this process are largely unknown. Given the importance of epigenetic remodeling in aging, here we analyzed the relationship between responsiveness to influenza vaccination and DNA methylation profiles in healthy subjects of different ages. Peripheral blood mononuclear cells were collected from 44 subjects (age range: 19 – 90 years old) immediately before influenza vaccination. Subjects were subsequently classified as responders or non-responders according to hemagglutination inhibition assay 4-6 weeks after the vaccination. Baseline whole genome DNA methylation in peripheral blood mononuclear cells was analyzed using the Illumina® Infinium 450k microarray. Differential methylation analysis between the two groups (responders and non-responders) was performed through an analysis of variance, correcting for age, sex and batch. We identified 83 CpG sites having a nominal p-value < 0.001 and absolute difference in DNA methylation of at least 0.05 between the two groups. For some CpG sites, we observed age-dependent decrease or increase in methylation, which in some cases was specific for the responders and non-responders groups. Finally, we divided the cohort in two subgroups including younger (age < 50) and older (age ≥ 50) subjects and compared DNA methylation between responders and non-responders, correcting for sex and batch in each subgroup. We identified 142 differentially methylated CpG sites in the young subgroup and 305 in the old subgroup, suggesting a larger epigenetic remodeling at older ages. Interestingly, some of the differentially methylated probes mapped in genes involved in immunosenescence (*CD40*) and in innate immunity responses (*CXCL16*, *ULK1*, *BCL11B*, *BTC*). In conclusion, the analysis of epigenetic landscape can shed light on the biological basis of vaccine responsiveness during aging, possibly providing new appropriate biomarkers of this process.

## Keywords

DNA methylation; Influenza vaccination; Aging; Immunosenescence

## 1. Introduction

Influenza is an important public health challenge in our countries, with yearly epidemics responsible of significant mortality, morbidity and loss of productivity (Paules and Subbarao, 2017). Specific populations such as very young children, individuals aged 65 years and older, or subjects with pre-existent conditions (immunocompromised states, cardiovascular or cerebrovascular diseases, diabetes, chronic respiratory failure, pregnancy) are particularly vulnerable to this infection and at greater risk for complications.

Vaccination is the most effective method to prevent influenza infection. Annual vaccination with an injectable trivalent inactivated vaccine is recommended, especially for individuals aged 65 years or older. However, the protection delivered by these vaccines is incomplete. Rates of protective immune response to vaccination are frequently low in vaccinated subjects, with worsened responses in older adults (Jefferson et al., 2010; Osterholm et al., 2012).

Part of the poor vaccine efficacy in the elderly is due to immunosenescence (Haralambieva et al., 2015; Kennedy et al., 2016; Targonski et al., 2007) but molecular pathways associated with impaired vaccine responses remain incompletely understood. Identification of the mechanisms associated with the development of a protective immunity is of central importance in vaccinology, in order to improve our capacity to predict response to vaccination or develop potential interventions to improve the immune responses.

To date, several studies have been conducted to identify genome-wide changes in transcriptional profiles that correlate with clinical response to influenza vaccination (Bucasas et al., 2011; Nakaya et al., 2011, 2015; Obermoser et al., 2013; Thakar et al., 2015; Tsang et al., 2014; Zhu et al., 2010), by assessing genome-wide gene expression with microarrays before and/or after vaccination of subjects. These molecular signatures, associated with better antibody responses, were frequently enriched in immune pathways, especially with type I interferon signaling, antigen presentation pathways or B-cell proliferation. Thakar *et al.* identified a dysregulation in this gene signature in older adults, specially in frail subjects who were non-responders to vaccination (Thakar et al., 2015). Other large-scale profiling studies have tried to identify further relevant biomarkers that could predict vaccine response: in this attempt, Furman *et al.* identified nine immunological baseline predictors of protective immunity, with two of these variables involved in apoptosis (Furman et al., 2013). Finally, models integrating and combining transcriptomic data with additional data types to predict response to vaccination have been developed recently (Tsang et al., 2014; Zimmermann et al., 2017).

While transcriptomic data have been deeply studied in this field, few reports have been published regarding epigenetic aspects. DNA methylation has an important role in several biological processes, especially in aging (Sen et al., 2016), and is therefore an interesting candidate to be investigated. Furthermore, DNA methylation measures tend to be more stable than transcriptomic data within the short period (days-weeks) and are more reproducible from a technical point of view. Lu *et al.* discovered two relevant epigenetic variations in poor-responders to the vaccine directed against Hepatitis B virus (Lu et al., 2014). Concerning influenza vaccine, one recent study has identified numerous CpG sites showing associations with gene expression and other ones associated with the induction of the humoral immune response (Zimmermann et al., 2016).

To complete these findings and evaluate the effect of age on vaccination response, here we investigated baseline (that is, immediately before vaccination) genome-wide DNA methylation in peripheral blood mononuclear cells (PBMC) from 44 healthy donors, ranging from 19 to 90 years, who received influenza vaccination and were classified as responders and non-responders according to hemagglutination inhibition assays (HIA) after 28 days.

## **2. Materials and Methods**

### **2.1 Participants**

Participants (age range: 19 – 90 years) were recruited at the University of Miami Miller School of Medicine. Experiments were conducted using peripheral blood. Enrolled participants received the influenza vaccine in the pandemic season 2009 and in the season 2010-2011. Participants enrolled in the pandemic 2009 season received the subunit vaccine containing the A/California/7/2009 (H1N1) strain, whereas those enrolled in the 2010-2011 season received the Trivalent Inactivated influenza vaccine containing the following viral strains: A/California/7/2009 (H1N1), A/Perth/16/2009 (H3N2), B/Brisbane/60/2008. Whole blood samples were collected immediately before vaccination. PBMC were collected using Vacutainer CPT tubes (BD 362761). Cells were washed and cryopreserved. Appropriate signed informed consent was obtained from each subject prior to enrollment. The study was approved with IRB protocol #20070481. Each participant was asked questions regarding demographics, health behaviors, presence of symptoms associated with inflammatory conditions or respiratory infections at the time of enrollment. No one reported subclinical inflammatory conditions and/or had respiratory tract infections at the time of enrollment, nor was on any anti-inflammatory treatment or on medications known to alter the immune response. Participants were excluded if they had diseases known to alter the immune response.

### **2.2 Assessment of response to vaccination**

Immunogenicity of influenza vaccine in subjects was assessed by hemagglutination inhibition assays (HIA). For this purpose, blood samples were collected immediately before vaccination (baseline) and 4-6 weeks after to evaluate the *in vivo* response and identify responders and non-responders. Responders had at least a 4-fold increase in the reciprocal of the titers in response to the whole vaccine, which was the A/California/7/2009 (H1N1) strain in the 2009 season and the Trivalent Inactivated Influenza vaccine (containing A/California/7/2009 (H1N1), A/Perth/16/2009 (H3N2), B/Brisbane/60/2008) in the 2010-2011 season. Briefly, sera were pretreated with receptor destroying enzyme (RDE, Denke Seiken Co Ltd) for 20 hours at 37°C; in order to inactivate this enzyme, sera were then heated at 56°C for 60 minutes. Two-fold serial dilutions were done; 25 µL of diluted sera were incubated with an equal volume of 4 HA units of the 2009 vaccine or of the 2010-2011 vaccine, for 1 hour at room temperature and then 50 µL of a 1.25 % suspension of chicken red blood cells were added. After 2 hours of incubation at room temperature titers were determined.

### **2.3 Genome-wide DNA methylation analysis**

Genome-wide DNA methylation analysis was performed on PBMC collected immediately before vaccination (baseline) and cryopreserved. DNA methylation patterns are generally stable, highly reproducible and only slightly affected by freezing (Bulla et al., 2016). Total genomic DNA was extracted from PBMC using the AllPrep DNA/RNA Mini kit (Qiagen), according to manufacturer's instructions. DNA concentrations were determined using NanoDrop spectrophotometer. DNA was bisulfite-converted using the EZ DNA Methylation Kit (Zymo Research Corporation®) and analyzed on the Infinium HumanMethylation450 BeadChip (Illumina®) following manufacturer's instructions (Bibikova et al., 2011). Arrays were scanned by HiScan (Illumina®) and signal intensities were extracted from *.idat* files

using the *minfi* Bioconductor package (Aryee et al., 2014). Data were normalized using the `preprocessQuantile` function of the package *minfi*. Probes on the X and Y chromosomes were removed, as well as probes associated to a SNP. Identification of CpG sites with differential methylation between responders and non-responders to influenza vaccination was performed through an analysis of variance (ANOVA) model, correcting for sex and batch. CpG sites differentially methylated between responders and non-responders were defined as having a nominal p-value inferior to 0.01 and an absolute difference between values of responders and non-responders of at least 0.05. Figures were generated using R.

## **2.4 Epigenetic age estimation**

DNAm age, also referred to as epigenetic age, was calculated as described by Horvath (Horvath 2013), using the online age calculator freely available at the website: <https://dnamage.genetics.ucla.edu>.

### 3. Results

#### 3.1 DNA methylation differences in responders and non-responders to influenza vaccination

Forty-four subjects were included in the present study: 23 were responders to influenza vaccine, while 21 were considered as non-responders according to HIA assay. Characteristics of the subjects are summarized in Table 1.

	N of subjects	Sex	N of subjects aged $\geq 50$ years	Age in the Young group (mean and sd)	Age in the Old group (mean and sd)
<b>Responders</b>	23	10M / 13F	11	28.4 (8.0)	64.1 (12.5)
<b>Non-responders</b>	21	10M / 11F	12	33.4 (6.6)	63.2 (7.4)

**Table 1.** Demographic characteristics of the subjects involved in the study.

We used the Illumina® Infinium 450k microarray to generate genome wide DNA methylation data from PBMC collected pre-vaccination (baseline). As a first step in our analysis, we compared DNA methylation between responders and non-responders to influenza vaccination by ANOVA, correcting for age, sex and batch. No CpG sites withstood Benjamini-Hochberg false discovery rate correction for multiple testing (significant threshold: q-value  $<0.05$ ). We then considered less stringent criteria (uncorrected p-value inferior to 0.01 and absolute difference between mean methylation values of responders and non-responders of at least 0.05). In this way, we identified 83 differentially methylated CpG sites between the two groups of subjects, mapping in 52 annotated genes (Tables 2 and 3; Supplementary File 1), that were able to distinguish responders and non-responders in a multidimensional scaling (MDS) plot (Figure 1A). When looking at the genomic localization of these probes, we found 3 genomic regions in which multiple adjacent CpG probes showed differential methylation between responders and non-responders: 3 probes mapped within the chr1:57110663-57111337 island associated to *PRKAA2* gene (Figure 1B), 2 mapped within the chr6:160209941-160212015 island associated to *TCPI/MRPL18* genes and 2 mapped in *TRAPPC9* gene, one in the chr8:141348840-141349195 island and one in the shore of the chr8:141467218-141467927 island. The presence of multiple differentially methylated probes within the same region indicates a general epigenetic remodeling of the genomic tract, which is likely to be biologically relevant (Bacalini et al., 2015; Wessely and Emes, 2012).

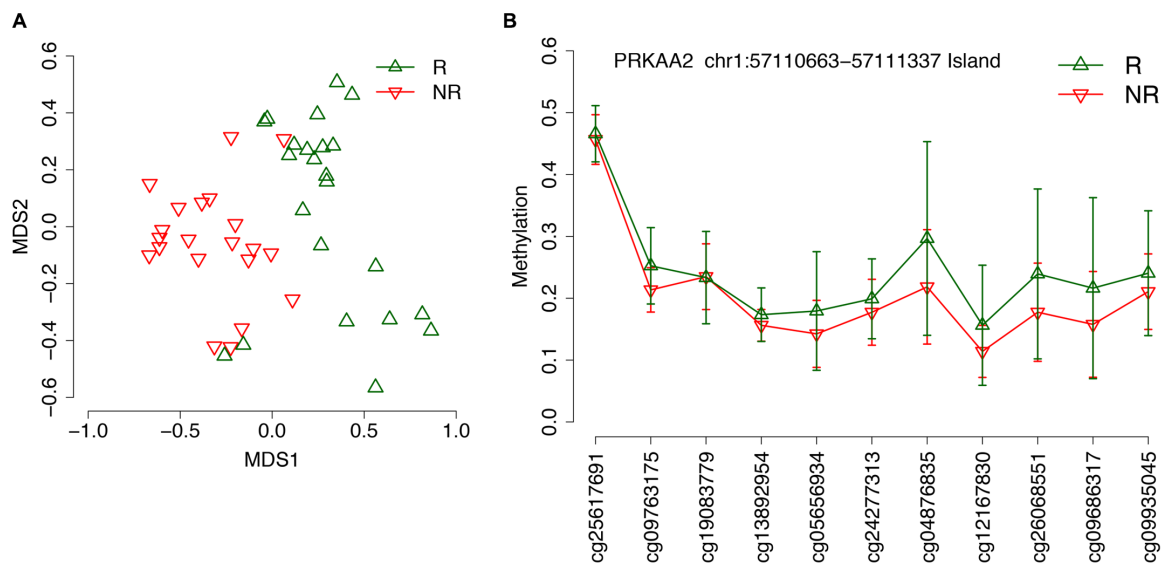
	Number of differentially methylated CpG sites between responders and non-responders
<b>All subjects</b>	83
<b>Young subjects (Age <math>&lt; 50</math> years)</b>	142
<b>Old subjects (Age <math>\geq 50</math> years)</b>	305

**Table 2.** Number of differentially methylated CpG sites between responders and non-responders to influenza vaccination, identified according to criteria described in Methods.



Illumina ID	Chromosome	UCSC_RefGene_Name	UCSC_CpG_Islands_Name	Relation_to_UCSC_CpG_Island	Uncorrected p-value	Difference between mean methylation values of R and NR
cg13235976	3		chr3:193858770-193859695	S_Shelf	3.04E-06	-0.0659
cg01281776	4		chr4:20253276-20256868	Island	0.00016	-0.0562
cg27559562	16	CLUAP1	chr16:3550773-3551274	N_Shore	0.00025	-0.0535
cg19492632	19	SLC7A9	chr19:33350719-33350932	N_Shelf	0.00028	0.0637
cg19497709	3	COPG	chr3:128997382-128997600	N_Shelf	0.00057	-0.0544
cg15219393	5	PPARGC1B	chr5:149109570-149111750	Island	0.00062	0.0582
cg05146089	5		chr5:926586-927401	Island	0.00068	-0.0507
cg22385827	2	C2orf67	chr2:211035478-211036637	Island	0.00072	0.0501
cg24845595	1	NTRK1; INSRR	chr1:156814881-156815792	N_Shore	0.00073	-0.0513
cg17177528	7		chr7:123672063-123673691	N_Shelf	0.00074	-0.0509

**Table 3.** Top 10 of the 83 differentially methylated CpG sites between responders (R) and non-responders (NR) to influenza vaccination.

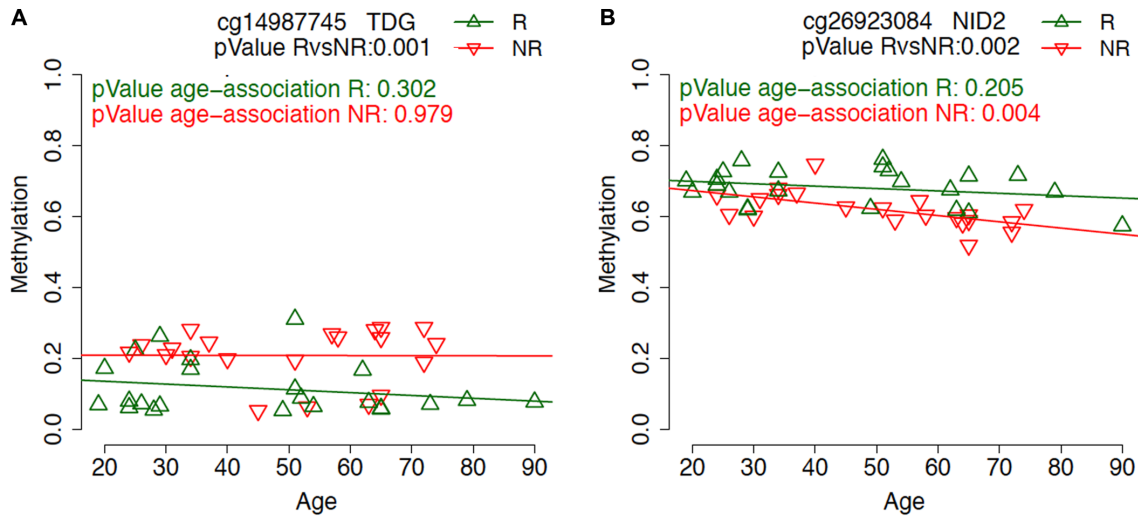


**Figure 1:** Comparison of DNA methylation profiles between responders (R) and non-responders (NR) to influenza vaccination. (A) Multidimensional scaling plots of DNA methylation data of 83 differentially methylated CpG sites between R (green) and NR (red). (B) Line plot of mean methylation values +/- standard deviation in R (green) and NR (red) for each CpG site mapping in the CpG island chr1:57110663-57111337 (*PRKAA2* gene).

### 3.2 Age-associated epigenetic determinants of vaccination response

In order to assess the effect of age on vaccination response, for each of the CpG probes identified as differentially methylated in the previous analysis we calculated the association with age, considering separately responders and non-responders to influenza vaccine

(Supplementary Files 1 and 2). For some CpG probes, such as cg14987745 in *TDG* gene, methylation differences between responders and non-responders were not affected by age (Figure 2A). On the contrary, others CpG sites showed age-association specifically in one of the two groups. For example, the probe cg26923084 in the gene *NID2* showed an age-dependent decrease in methylation only in non-responders, with larger DNA methylation differences after the age of 50 years old (Figure 2B).



**Figure 2:** Scatter plots of methylation values (y-axis) according to age of the subjects (x-axis) for the CpG probes in *TDG* (A) and *NID2* (B) genes. Values indicate the pValue of the association between methylation and age in responders (R) (green) and non-responders (NR) (red) subjects.

To better identify the epigenetic determinants of vaccine responsiveness that are dependent on the age of the subjects, we divided the cohort in two groups including younger (age < 50) and older (age ≥ 50) subjects. Within each subgroup, we compared DNA methylation between responders and non-responders, correcting for sex and batch. Also in this case, no CpG site survived the multiple test correction. Using the same selection criteria described above, in the Young group we identified 142 differentially methylated CpG sites, mapping in 98 annotated genes (Tables 2 and 4; Supplementary Files 3 and 4). Four loci had multiple differentially methylated probes: *C5orf33* (2 CpG probes in the chr5:36241003-36242664 island), *HCG4P6* (3 CpG probes in the chr6:29894140-29895117 island and its shore), *TRAF3* (2 CpG probes in the body of the gene) and *RPTOR* (4 CpG probes in the body of the gene, in the chr17:78863569-78863813 island and in the adjacent shore). In the Old subgroup, 305 CpG probes turned out to be differentially methylated, mapping in 206 genes (Tables 2 and 5; Supplementary Files 3 and 5). Eight loci contained multiple differentially methylated probes between old responders and non-responders: *KIF5C* (2 CpG probes, one in the chr2:149632682-149633882 island and one in the shore of chr2:149645536-149645834 island), *COL7A1* (2 CpG probes in the chr3:48631882-48632901 island), *MIR886* (2 CpG probes in the chr3:48631882-48632901 island), *TRAPPC9* (2 CpG probes, one in the chr8:141348840-141349195 island and one in the shore of chr8:141467218-141467927 island), *NKX2-3* (2 CpG probes in the chr10:101294443-101297263 island), *MCF2L* (2 CpG

probes, one in the chr13:113687421-113687828 island and one in the shore of the chr13:113714784-113715160 island), *MYOD* (2 CpG probes, one in the shore of chr17:30822917-30823210 island and one in the chr17:30845903-30846702 island) and *RGL3* (2 CpG probes, one in the chr19:11531278-11531590 island and one in its shore). No CpG probes were identified as differentially methylated both in the Young and in the Old group; however, when considering the genes in which the CpG probes map, *COL7A1* and *PAPLN* turned out to be present in both the comparisons. For both the genes, the CpG probes identified as differentially methylated in the two comparisons mapped in near positions along the chromosome (within 1000 bp) and showed consistent age-dependent trends (for both the genes, and age-related hypomethylation specifically in the non-responders group), suggesting a general epigenetic remodeling of the two genomic regions.

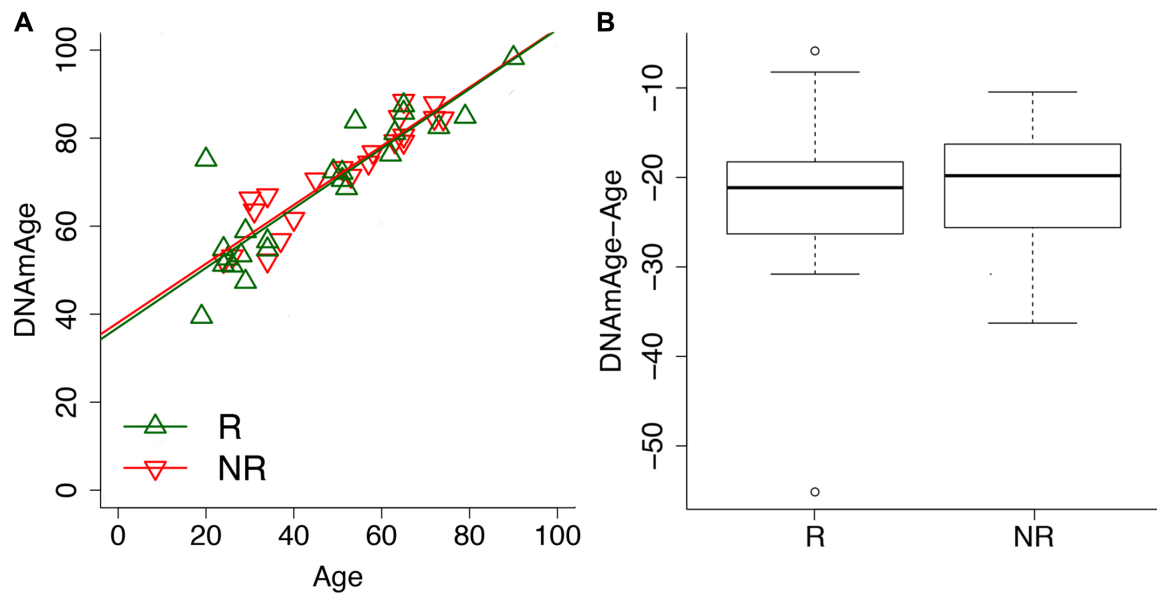
Illumina ID	Chromosome	UCSC_RefGene_Name	UCSC_CpG_Islands_Name	Relation_to_UCSC_CpG_Island	Uncorrected p-value	Difference between mean methylation values of R and NR
cg21690945	4		chr4:170695989-170696274	S_Shelf	0.00011	-0.0520
cg10420952	7	TWIST1	chr7:19156050-19158042	Island	0.00012	-0.0591
cg25622597	3	RHOA; TCTA	chr3:49448861-49449965	Island	0.00024	0.0626
cg10033725	7	HEATR2	chr7:821689-822634	S_Shelf	0.00029	-0.0662
cg24233211	6				0.00031	-0.1159
cg07681156	19	ZNF581	chr19:56154791-56155117	Island	0.00038	-0.0551
cg14562060	6	TCP11	chr6:35108801-35109499	Island	0.00040	-0.0553
cg05696877	1	IFI44L			0.00046	0.0573
cg23696891	6	TNFAIP3	chr6:138187825-138189141	Island	0.00052	-0.0631
cg17014647	6	ATXN1			0.00067	0.0589

**Table 4.** Top 10 of the 142 differentially methylated CpG sites between responders (R) and non-responders (NR) to influenza vaccination (young subjects with age < 50 years).

Illumina ID	Chromosome	UCSC_RefGene_Name	UCSC_CpG_Islands_Name	Relation_to_UCSC_CpG_Island	Uncorrected p-value	Difference between mean methylation values of R and NR
cg14273502	12		chr12:113013099-113013529	Island	6.99E-05	-0.0575
cg11644052	2	NFU1	chr2:69664291-69664816	S_Shore	0.00012	-0.0567
cg26923084	14	NID2	chr14:52534581-52536722	S_Shore	0.00013	-0.0899
cg23301925	14				0.00021	-0.0501
cg22256027	4	MSX1	chr4:4864456-4864834	N_Shore	0.00024	0.0586
cg13617280	12	SLC15A4; MGC16384			0.00029	0.0825
cg24927800	2	DES	chr2:220283200-220283750	Island	0.00031	-0.0710
cg16956133	2		chr2:863930-865091	Island	0.00035	-0.0861
cg21126344	17	SOX9	chr17:70116274-70119998	Island	0.00036	-0.0816
cg10857729	8		chr8:976015-976416	N_Shelf	0.00043	0.0622

**Table 5.** Top 10 of the 305 differentially methylated CpG sites between responders (R) and non-responders (NR) to influenza vaccination (old subjects with age  $\geq 50$  years).

Finally, we investigated if the responsiveness to vaccination was associated to the epigenetic age of the subjects. In the last few years, DNA methylation-based biomarkers have gained particular relevance in the field of aging research and have been proposed as proxy of the biological age of an individual. The most popular epigenetic age estimator is Horvath's epigenetic clock (Horvath, 2013) which has been shown to correlate with mortality (Chen et al., 2016; Christiansen et al., 2016; Marioni et al., 2015a; Perna et al., 2016), physical and cognitive fitness (Marioni et al., 2015b) and to detect a biological age-acceleration in frail subjects (Breitling et al., 2016). Frailty is of particular importance from our perspective, since it has been observed that this status impacts susceptibility to influenza and responsiveness to influenza vaccine (Yao et al., 2011). Here we used Horvath's epigenetic clock to estimate the DNA methylation age (DNAMAge) in our cohort. Then for each individual, we calculated the delta between chronological age and DNAMAge, in order to detect deceleration (the subject is epigenetically younger than his/her chronological age) or acceleration effects (the subject is epigenetically older than his/her chronological age). No significant differences in delta values were observed between responders and non-responders, nor when we divided the cohort in the young and old subgroups (Figure 3).



**Figure 3:** Estimation of Horvath's epigenetic clock in responders (R) and non-responders (NR) to influenza vaccination. (A) Scatter plot of DNAmAge (y axis) versus chronological age (x axis) in R and NR to influenza vaccination. (B) Boxplot of differences between DNAmAge and chronological age in R and NR.

#### 4. Discussion

Understanding mechanisms associated with impaired vaccine responses is an important goal and this is particularly true in the context of aging. Age-associated alterations in immune responses, named immunosenescence, are in part responsible of the poor response after vaccination in elderly (Goronzy and Weyand, 2013). Role of DNA methylation in this phenomenon has been investigated in subjects aged 50 – 74 years old vaccinated against influenza. The authors identified multiples CpG sites associated with age and with strong correlations with immunosenescence markers (Kennedy et al., 2016), highlighting the possible implication of epigenetic regulation as an active mechanism that shapes the immune response during aging.

In this study, we analyzed baseline whole genome DNA methylation profiles of 44 subjects ranging 19-90 years and receiving influenza vaccination. To the best of our knowledge, this is the first study to investigate methylation data in vaccinated subjects with such a wide range of ages, making it possible to identify age-related determinants of influenza vaccination response. Accordingly, we identified 83 differentially methylated probes when correcting for the age of the subjects, 142 in the young subgroup and 305 in the old subgroup. This suggests that DNA methylation differences between responders and non-responders are larger at older ages compared to younger ones. Non-responders did not show epigenetic age acceleration effects compared to responders, according to Horvath's epigenetic clock.

To our knowledge, none of the CpG sites we identified was previously found differentially methylated in studies investigating DNA methylation profiles in human vaccinology, and no significant enrichment in KEGG pathways was observed (data not shown). However, it is interesting that we identified differentially methylated probes mapping in genes that are

involved in immunosenescence (CD40) (Metcalf et al., 2015; Toapanta and Ross, 2009) and in innate immunity responses during viral infection: CXCL16 (Piqueras et al., 2006), ULK1 (Prantner et al., 2017), BCL11B (Yu et al., 2015), BTC (Al-Yahya et al., 2015).

Senescence of the adaptive compartment of immunity, which is involved in the classic immunological memory (Crotty and Ahmed, 2004), has been widely studied and most of the cellular markers studied in the report published by Kennedy *et al.* are related to B or T lymphocytes (Kennedy et al., 2016). However, during last years, evidence has emerged that also innate immune cells have a sort of memory, called “trained immunity”. This phenomenon could be a contributor to the protective effects of vaccination, independently of T and B cells, mostly observed in bacteriological vaccines, such as BCG (Garly et al., 2003; Kleinnijenhuis et al., 2012; Tribouley et al., 1978; van 't Wout et al., 1992). Whereas immunological memory associated to adaptive immunity relies upon genetic recombination and mutations for the development of antigen specific receptors, mechanistic studies have brought evidence that trained immunity does not involve these types of permanent genetic changes but is largely based on epigenetic remodeling in innate immune cells (such as myeloid cells (monocytes, macrophages), natural killer cells or innate lymphoid cells) (Netea et al., 2016). The epigenetic shift, mostly based on histone modifications but also on microRNAs and DNA methylation changes, follows modifications in cellular metabolism driven by the antigenic stimulation (Arts et al., 2016) and can shape the cellular response and remains over time. Identification of differentially methylated CpG sites between responders and non-responders after vaccination is an important step to gain knowledge on these processes, for the identification of genes that could be potential targets for functional analysis and for the development of new strategies of vaccination.

In this study we analyzed total PBMC samples, like in most published system vaccinology studies which used whole blood or PBMC. One important advantage of this approach is that blood and PBMC can be easily obtained from individuals, but an important limit is the presence multiple diverse cell subsets. Indeed, the heterogeneity in cell populations in the analysis can possibly hide differences in DNA methylation patterns in individual cell types. Hoek *et al.* previously investigated the transcriptomic and proteomic profiles from PBMC and from sorted immune cells after influenza vaccination (T cells, B cells, natural killer cells, myeloid dendritic cells, monocytes and neutrophils). They described significantly different RNA and protein expression profiles between the groups (Hoek et al., 2015).

Another possible limitation in our work is the lack of data regarding pre-existing immune status of the subjects that could affect vaccine response. It is known that older subjects are frequently non-responders to vaccine and that a 4-fold increase in HIA titers is more frequently found in young subjects as compared to old ones: in a meta-analysis, Goodwin *et al.* generated an adjusted odds-ratio of responses in elderly vs young adults of 0.24 to 0.59 in terms of seroconversion and seroprotection against 3 different influenza antigens (H1N1, H3N2 and B) (Goodwin et al., 2006). Weakened serological response to vaccination in elderly subjects are related to the effect of immunosenescence, but previous exposures to influenza antigens (*via* vaccinations or infections) have also an influence on the response to the new vaccination. In general, subjects with repeated previous influenza vaccinations and influenza specific antibodies at baseline have a reduced humoral response and so lower post-vaccination

titers (Bucasas et al., 2011; Gulati et al., 2005; Huang et al., 2017; Ng et al., 2013; Sasaki et al., 2008; Thakar et al., 2015; Tsang et al., 2014). However, it has also recently been demonstrated in a study conducted on 136 young and 122 elderly individuals that young individuals respond better than elderly individuals to the first vaccination, but after subsequent vaccinations the difference in response between young and elderly individuals declines rapidly, suggesting that different prior vaccination history and/or infection histories must also be taken in consideration when influenza vaccination is examined (Mosterin Höpping et al., 2016).

In this study, we evaluated the association of baseline DNA methylation patterns and response to the vaccination at different ages. Future studies should systematically evaluate the epigenetic response to influenza vaccination in subjects of different ages, analyzing DNA methylation and gene expression at different time points post-vaccination and correlating changes in epigenetic patterns with the antibody response. Although it has been demonstrated that vaccination does not profoundly alter DNA methylation profiles in PBMC (Zimmermann et al., 2016), further studies are necessary in this field, considering the impact of the age of the individuals and analyzing different cell types separately. Furthermore, DNA methylation signatures should be explored and validated in other independent datasets considering other influenza strains.

## **5. Conclusion**

In conclusion, in this paper we identified possible age-related DNA methylation contributors to vaccine responsiveness. Further studies on larger independent cohorts are needed to define epigenetic biomarkers that can predict the vaccine efficacy. These models could be used to develop novel strategies to achieve optimal protective immune responses, specially in the elderly.

**Conflicts of interest:** none

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## Figure Legends

**Figure 1:** Comparison of DNA methylation profiles between responders (R) and non-responders (NR) to influenza vaccination. (A) Multidimensional scaling plots of DNA methylation data of 83 differentially methylated CpG sites between R (green) and NR (red). (B) Line plot of mean methylation values  $\pm$  standard deviation in R (green) and NR (red) for each CpG site mapping in the CpG island chr1:57110663-57111337 (*PRKAA2* gene).

**Figure 2:** Scatter plots of methylation values (y-axis) according to age of the subjects (x-axis) for the CpG probes in *TDG* (A) and *NID2* (B) genes. Values indicate the pValue of the association between methylation and age in responders (R) (green) and non-responders (NR) (red) subjects.

**Figure 3:** Estimation of Horvath's epigenetic clock in responders (R) and non-responders (NR) to influenza vaccination. (A) Scatter plot of DNAmAge (y axis) versus chronological age (x axis) in R and NR to influenza vaccination. (B) Boxplot of differences between DNAmAge and chronological age in R and NR.

## Supplementary material

**Supplementary File 1:** List of the differentially methylated CpG probes between responders (R) and non-responders (NR).

**Supplementary File 2:** Scatter plots of methylation values (y-axis) according to age of the subjects (x-axis) for the 83 CpG probes identified as differentially methylated between responders (R) (green) and non-responders (NR) (red) subjects. Values indicate the p-values of the association between methylation and age in responders (R) (green) and non-responders (NR) (red) subjects.

**Supplementary File 3:** Lists of the differentially methylated CpG probes between responders (R) and non-responders (NR) in young and old subjects.

**Supplementary File 4:** Scatter plots of methylation values (y-axis) according to age of the subjects (x-axis) for the 142 CpG probes identified as differentially methylated between in responders (R) (green) and non-responders (NR) (red) subjects in the young subgroup. Values indicate the p-values of the association between methylation and age in responders (R) (green) and non-responders (NR) (red) subjects.

**Supplementary File 5:** Scatter plots of methylation values (y-axis) according to age of the subjects (x-axis) for the 305 CpG probes identified as differentially methylated between in responders (R) (green) and non-responders (NR) (red) subjects in the old subgroup. Values indicate the p-values of the association between methylation and age in responders (R) (green) and non-responders (NR) (red) subjects.